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(54) Title: MONOCLONAL ANTIBODIES TO RENAL CELL CARCINOMA

(57) Abstract

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against an antigen or mixture of antibody and their use in theraphy and diagnosis of RCC are also described. cells and to varying extent on other benign and malignant tumors and normal cells. Compositions containing antibody Antigens associated with renal cell carcinoma (RCC) are described. The antigens are present on renal carcinoma

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MONOCLONAL ANTIBODIES TO RENAL CELL CARCINOMA

Background of the Invention

particularly difficult to diagnose. See Oberling et Renal cell carcinoma (RCC) is a cancer which is treatment of these tumors in vivo. can also be useful for detection, localization and Radiolabeled Mabs specific for these antigens tests for tumor associated or tumor specific antireagents in diagnostic tumor pathology, and in serum Mabs recognizing such antigens can be useful in (1981) and Magnani et al., Cancer Res. 43:5489-5492. (1983); Wilson et al., Int. J. Cancer 28; 293-300 (1981). Wright et al., Cancer Res. 43, 5509-5516 et al. Proc. Nat'l. Acad. Sci., 78: 3199-3203 ficity for these tumor antigens. See e.g., Colcher monoclonal antibodies (Mabs) having a high specipresent only on malignant tumors and to isolate efforts have been made to identify antigens that are (3797) (375) 495-497 (1975), Kohler and Milstein, nique for production of nonspecific antibody by Since the introduction of the hybridoma tech-

al. <u>Nature</u> 186, 402-403 (1960) and Holthafer et al., Lab. Invest. 49, 319-326 (1983). Mabs specific to antigens which are pre-dominantly present only on RCC tumor cells would be extremely useful in the diag-

nosis and treatment of this disease.

KC 38 TE

Summary of the Invention

associated with renal cell carcinoma (RCC) and to

and are characterized by distinct patterns of tissue are designated G 250, RC 38, RC 3, RC 69 and RC 154

visceral glomerular epithelial cells at the capillary

of the sweat glands, and on the sinuses of the lymph the colon, on the sinuses of the liver, on the acini the crypts and villi of the jejunum, on the crypts of also present on the mucous cells of the stomach, on

with 8 out of 13 RCC metastases. No reaction was

(except for the epithelium of the bile ductand

and absent on normal adult fetal kidney tissue.

body reacted with 46 out of 47 primary RCC tested and primary and metastatic RCC tumors. The RC 38 anti-

Ag G 250 is absent from all other normal adult tissue

Antigen RC 38 is an antigen which is present on

Antigen (Ag) G 250 is an antigen present on RCC

seen with 179 tumors of various origin.

stomach) and most malignancies.

distribution as elaborated below.

If is also present on the glomerular visceral

Further, RC 38 is present on the differentiating loop on tissue sections of normal adult kidney. tubules up to the thin descending part of Henle's epithelium and the epithelial cells of the proximal

therapeutic compositions useful in these methods. and method of treatment of RCC and diagnostic and The invention also pertains to methods of diagnosis monoclonal antibodies reactive with the antigens.

The RCC associated antigens of this invention This invention pertains to several new antigens

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metanephros. tubules connected to these regions of the fetal loop stage and on the most proximal part of the

marginally present on the parietal epithelial cells tissue sections of normal adult kidneys. RC 3 is and the thick descending part of Henle's loop in Both RC 3 and RC 69 are present on proximal tubules present on primary RCC but absent on metastic RCC. marginally present on metastic RCC. Antigen RC 69 is Antigen RC 3 is present on primary RCC and

tissues which were tested. RC 3 and RC 69 were not present on any non-renal metanephros at the middle limb of the S-shaped stage. outgoing tubule. RC 69 is present on the fetal of Bowman's capsule in the region adjoining the

byros, on the ducts of adult breast glands and on the It is present on the fetal metaneadult kidney. tubules and small collecting ducts of the normal on the proximal tubular epithelium and on the distal absent on metastic RCC. RC 154 is marginally present

Antigen RC 154 is present on primary RCC and

RC 38, RC 3, RC 69 and RC 154 can be used in the Antibodies reactive with the RCC antigens G 250, follicles of the adult thyroid gland.

of RCC adjacent to uninvolved kidney tissue with Mab Figure 1 (Left) shows immunoperoxidase/(DAB) staining

Brief Description of the Drawings

diagnosis and treatment of RCC.

negative. tumor cells (bottom), uninvolved kidney tissue G 250. (Magnification 64x.) Diffuse staining of RCC

staining of RCC tumor cells is visible. (Magnification 512x.) Diffuse, membranous Right: GAM_FITC staining of primary RCC with

the staining is visible in the inset. canuliculi is visible. The cytoplasmatic aspect of cation inset 880x) Cytoplasmatic staining of bile normal liver with G 250 (magnification 160x; Magnifi-Figure 2. shows immunoperoxidase/DAB staining of

Figure 3.

liver is visible. In the melanoma bearing mouse (fig 3c), only the 3a the injection point in the tail is also visible. are visible 2 and 20 hours after injection. In fig In figure 3a, b the RCC tumor as well as the liver intraveneous injection of 1.5 ug^{99m} Tc labeled G 250. jection, 50,000 count images. Mice were given an melanoma tumor bearing mouse, 20 hours after in-Fig 3c: Radioimmunoscintigraphy of a after injection (fig 3b), 100,000 count images per tumor bearing mouse, 2 hours (fig 3a), and 20 hours Fig 3a, b: Radioimmunoscintigraphy of an RCC

reaction of malignant tumors with G 250. (black: Figure 4 is a bar histogram indicating staining

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more than 50% of tumor cells stained; striped: more than 1% and less than 50% of tumor cells stained; than 1% and less than 1% of tumor cells stained; white: no tumor cells stained; ca = carcinoma). Numbers no tumor cells stained; ca = carcinoma). Numbers between brackets are total numbers of tumors tested. These represent mostly primary tumors. The metalsylumors (one with less than 1% tumor (one with less than 1% tumor cells positive, eight negative), four pulmonary tumors (all negative) and four colonic tumors (one with more than 50% positive tumor cells, onw with less than 1% tumor cells positive). Percentages in bars represent percentage of tumors with staining characteristics corresponding of tumors with staining characteristics corresponding of tumors with staining characteristics corresponding to bar color.

Detailed Description of the Invention

The antibodies of this invention react with antigens present on RCC. The antigens are designated G 250, RC 38, RC 3, RC 69 and RC 154. In brief, RC 38 reacts with 95% of primary and 67% of meta-static RCC and did not react with other tumors tested (but did react with some normal tissue). G 250 reacts with primary and metastatic RCC but not with normal tissue. RC 4 reacted with RCC and a wide normal tissue. RC 4 reacted with RCC and a wide normal tissue.

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The presence of Ag G 250 was determined by the staining reaction of the antigen with Mab G 250. G 250 is present in a high percentage of cells in most RCC tumors and is absent from the cells of

The fact that G 250 stains a variety of other tumors, although with low incidence, makes it less suitable for establishing a differential diagnosis of RCC. The strong staining of cell membranes in the majority of RCC tumor cells in most RCC suggests that this Mab can be useful for tumor scintigraphy. In radioimmunoscintigraphic experiments, RCC tumors from radioimmunoscintigraphic experiments, RCC tumors from

frequently in colonic carcinomas. negative. G 250 antigen is expressed relatively these cases the corresponding normal tissues were in carcinomas but occasionally also in sarcomas. determinant was also found in nonRCC tumors, mainly activation of a cellular oncogene product. The G 250 possibly due to a common initiating event such as synthesis is inherently related to tumor development, primary RCC suggest that induction of G 250 antigen adenoma and the general occurrence of this antigen in The G 250 appearance in renal histochemistry. denced by the data obtained in ELISA and immunoto be absent from the normal adult kidney as evitwo renal adenomas tested. The G 250 antigen appears benign tumors and premalignant lesions including the Expression of G 250 antigen was found in a few

normal kidneys. With respect to the normal tissues tested G 250 is present on normal bile duct epither lium, mucous cells in the stomach and is marginally rescrion was found with other adult tissues tested nor with any fetal tissue tested.

The stop of C 250 appials to the stop was found in a few portion with any fetal tissue tested.

a cell line with moderate expression of G 250 antigen (as estimated from the immunofluorescence data) were visualized. Clear tumor resolution was obtained with a diameter ranging from 5-7mm without using subtraction techniques. Tumors were also visualized with labeled G 250 in tumor bearing subtraction is present in only a few normal tissues suggests that antibodies specific to G 250 antigen sre useful for RCC diagnosis.

The RC Series

Antigens RC 3, RC 69, RC 154 and RC 38 were also discovered on the cells of renal cell carcinoma tumors. These antigens were discovered by their reaction with a series of monoclonal antigen with Mab RC 3, RC 69 antigen with Mab RC 154 and RC 38 antigen with Mab RC 154 and RC 38 antigen with Mab RC 38. These monoclonal antibodies and the cell lines which produced them are specific antigen with Mab RC 38. These monoclonal antibodies and the cell lines which produced them are specific antigen with Mab RC 38. These monoclonal antibodies and the cell lines which produced them are specific antigen with Mab RC 38. These monoclonal antibodies and the cell lines which produced them are specific embodiments of the monoclonal antibodies and cell

By their reactions with their respective Mabs, the expression of the antigens RC 3, RC 69, RC 154 and RC 38 was found to be variable both with respect to numbers of antigen expressed and to the percentage of antigen-positive RCC tumor cells. The combinations of antigen expression RC 3+/RC 69+/RC 154+ and tions of antigen expression RC 3+/RC 69+/RC 154+ and primary RCC whereas no tumors were found to have a

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154+ or RC 3-/RC 69-/154+, combination of antigen expression RC 3-/R C69+/RC

in RCC metastases while RC 3 was found on only 1 out RC 69 antigen and RC 154 antigen were not found

of 10 of the metastases tested.

The data indicates that RCC Cells with metasta-

sizing capacity have lost the antigens corresponding

antigen, while six others did not express RC 38 Of the renal adenomas only one expressed RC 38 to RC 3, RC 69 and RC 154.

are compatible with an origin from distal tubular primary RCC always express RC 38 antigen. tubules to RCC as proximal tubular epithelium and represent a transitional stage from normal proximal only suggest that a subset of these adenomas do not Our limited data on small renal adenomas Bnckley, A. No. 49) Geneva, UICC, 1980, Eds. Sufrin, G. and Renal Adenocarcinoma (UICC Technical Reporting Series See Bennington, "Histopathology of Renal Tumors" in: cm constituting the borderline between these lesions

considered to be the size of the lesion, diameter 3 distinguishing point between adenoma and carcinoma is

considered precursor lesions of RCC and the main stained in all sections. Renal adenomas are often surrounding proximal tubular epithelium was clearly

antigen. RC 3, RC 69 and RC 154 antigen were not

expressed in the two renal adenomas tested.

breast tumors, carcinomas of the gastrointestinal RC 38 antigen is not present on tumor cells of •muiladiide

tract or angiosarcomas. In most tumors, RC 38 antigen is present on the endothelium of small vessels whereas it is absent in capillary endothelium of normal tissues except in the liver and lymph nodes. Therefore RC 38 might be useful in studies on angiogenesis or vascularisation in human tumors. Combining the data of tables 4 and 8 it is seen

that tumor cells of 46 out of 47 primary RCC and 8 out of 13 metastatic RCC were stained with RC 38. Also, RC 38 did not stain tumor cells of a wide variety of other tumors that included 12 clear cell tumors of different origins. These data indicate that RC 38 is useful for diagnostic purposes.

Immunoassay for RCC antigen Antibodies against the enumerated RCC antigens

can be used to detect RCC antigen in samples of bodily fluids (e.g. serum or plasma). For example, serological tests for circulating antigen may have diagnostic or prognostic value. Immunoassays for detection of RCC antigens can be performed in any of the standard formats such as competitive or immuno-metric formats.

Immunohistochemical Staining Human tissue specimens (e.g. biopsy samples) can

be tested for the presence of the RCC antigens by immunohistochemical techniques such as immunoperoxidase dase staining. As an alternative to immunoperoxidase staining immunofluorescent techniques can be used to

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examine human tissue specimens with the anti-RCC antibodies. In a typical protocol, slides containing cryostat section of frozen, unfixed tissue biopsy incubated with the anti-RCC antibody in a humidified anti-RCC antibody in a humidified anti-RCC antibody. For example, if a murine anti-RCC antibody is used the second antibody can be an anti-mouse antibody. The second antibody is labeled anti-mouse antibody. The second antibody is labeled anti-mouse antibody. The second antibody is labeled by fluorescent light microscopy.

Immunoscintigraphy

An immunoscintigraphic image of RCC in vivo can be obtained by administering to a person suspected of having RCC, labeled antibody (or a mixture labeled of antibodies) against an RCC antigen and allowing sufficient time for the antibody to accumulate at the antibody is then detected by an appropriate detecting antibody is then detected by an appropriate detecting device and the detected by an appropriate detecting image of the tumor. The constructed image can be used to localize and to assess the size of the tumor in vivo.

As immunoscintigraphic agents, antibodies

against antigens G250 and RC 38 are preferred.

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(i.e., "cocktails") can also be used. For example, a seminated tumor. As noted, mixtures of antibody and thus can abe used to localize primary and disantigens are expressed by primary and metastatic RCC

properties of both antibodies. an imaging composition having the advantages of the mixture of the antibody G 250 and RC 38 can provide

isotopically labeled intact antibody or antigen-For radioimmunoscintigraphy in humans, radio-

The preferred label for immunoscintigraphy is a Goldenberg in U.S. Patent 4,331,647. (1982) J. Biol. Response Modifiers 1, 121-136 and by are described by Goldenberg, D.M. and Deland, F.H. antibody fragments in tumor radioimaging techniques fragment may be used. The advantages of using monovalent Fab' fragment or the divalent F(ab') $_2$ binding fragments of anti-RCC antibody such as the

antibody or antibody fragment. For instance, the sciq (DTPA); any of these may be used to label the lating agent such as diethylene triamine pentacetic isotopes to proteins either directly or via a che-A variety of methods exist for attaching the radio-125 Lodine, 131 Lodine, 99m Technetium or 111 Lndium. radioimaging techniques including 123 Iodine, isotopes conventionally employed in in vivo tumor gamma camera). Examples of gamma emitting radiowith a conventional photoscanning device (such as a gamma-emitting radioisotope which can be detected

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magnetic resonance (MMR) properties of tissues pounds containing manganese) can also be employed. Monoclonal antibodies may be used to deliver the

ceptable vehicle.
Imaging based on the detection of nuclear

provided in kits for radioimmunoscintigraphy in humans. Preferably, such a kit includes either antibody the monovalent fragment Fab', the bivalent fragments (e.g. G250 and RC 38). In general, the labeling procedure will be prepared by the clinician. The antibodies or fragments can be provided with a preattached chelator (e.g. DTPA) for provided with a preattached chelator (e.g. DTPA) for provided with a preattached chelator (e.g. DTPA) for prepared for injection in a physiologically action in a physiological physio

The immunoscintigraphic composition is injected into the patient intravenously, intra-arterially or intraperitioneally. The amount of radioactivity infraperitioneally. The amount of radioactivity injected should be sufficient for detection by a standard gamma camera after the labeled antibody has distributed through the tissues of the body.

The anti-RCC antibodies of this invention may be

antibody may be labeled with Na[l25] by the chlor-amine—T method. See Hunter, W. M. and Greenwood, F. C. (1962), Nature 194, 495. Antibody may be directly labeled with 99m Technetium by the technique of Crockford et al., U.S. Patent No. 4,424,200, or it may be attached via a DTPA chelate as described by Hnatowich, U.S. Patent 4,479,930. In general, the antibody or antibody fragment is labeled to an antibody or antibody fragment is labeled to an appropriate specific activity (generally at least about 5 uCi/ug protein).

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paramagnetic substance to the RCC site and allow detection of tumor masses by MMR imaging.

Therapeutics

Tinking. tional techniques such as glutaraldehyde crossagents may be attached to the antibody by convensitizers such as boroncontaining organics. radiosensitizers such as misanidazole or neutron senradionuclides such as ^{ZII}Astatine and ^{I3I}Iodine, toxins such as the subunit of diphtheria toxin, include antibiotics, lectins such as ricin and abrin, analogs mercaptopurine and fluorouracil. Others folate, methotrexate, or the purine or pyrimidine RCC antibody are antimetabolites, such as the antior cytotoxic agents that may be linked to the anti-Among the various antiproliferative, antineoplastic given alone or as a carrier of an anti-tumor drug. patients afflicted with RCC. The antibody can be therapeutically effective (anti-tumor) amounts to targeted to RCC antigens can be administered in YbodijnA . also provide a basis for therapy of RCC. The RCC-associated antigens of this invention

Targeting Cytotoxic Cells Antibodies against the RCC antigens can be used

to target cytotoxic cells (e.g. human T cells, monocytes or NK cells). Cytotoxic cells can be attached to RCC via Fc receptors on the cells (which bind the Fc portion of an anti-RCC antibody) or via a

bridging antibody of dual specificty (i.e. an anti-body specific for the cytotoxic cell and for RCC).

by chemically coupling two antibodies of desired antibodies can also be produced as heteroantibodies targeting cytotoxic T cells to RCC. Bispecific line which produces T3/RCC bispecific antibody for antigen (preferably RC 38 or G250) to yield a cell with hybridoma producing antibody against the RCC hybridoma producing an anti-T3 antibody can be fused Harbor Symposium Quant. Biol. 1977: 41, 793. See, e.g. Immunol. Rev. 1979; Cold Spring specificity of the antibodies produced by the par-(quadroma) will produce hybrid antibody having a hybridoma producing anti-cytotoxic cell antibody fusion of a hybridoma producing anti-RCC antibody and cell to be targeted. For example, a cell formed by with a cell producing antibody against the cytotoxic can be produced by fusing an anti-RCC producing cells Dual or bi-specific antibodyes for targeting RCC

chemotherapy of RCC.

Class Switch Variants

Monoclonal antibodies against the RCC antigens

an adjunct to surgical therapy, radiation therapy, or

The cytotoxic cell can be targeted by allowing

targeting, the cells can be adminstered to the

the bispecific antibody to bind the cell.

specificity.

Therapy with targeted cells can be used as

G 250, RC 38, RC 3, RC 69 and RC 154 of different

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The invention is illustrated further by the allergic effect at the tumor site. variants may interact with mast cells to provide an M, D and E classes can also be produced. For receptors on effector cells. Antibody of the A, 2a, 2b and 3) can be produced which bind to different 82,8653. Antibodies of the various G subclasses (1, 1983:131 877; PNAS 1980 77, 2909 and PNAS 1985; e.g. J. Immunol., 1982:128, 1271; J. Immunol. tor class switching of hybridoma antibodies. rwwnuodjopnjiu cjases can be prepared by techniques

following examples.

EXYMBIE J

Production of the G 250 Hybridoma Cell Line

coated with 0.5% gelatin was also applied to identify used for the immunization procedure. A second filter coated with a mixture of 4 RCC cell homogenates not et al., Proc. Natl. Acad. Sci., 76, 1420-1424 (1979), agar was overlaid with nitrocellulose filters, Sharon cultured in soft agar. After 10 days of growth, the Kohler and Milstein, Nature, 256, 495-497 (1975), and with Sp2/0 myeloma cells essentially according to the spleen cells were isolated. These were fused the last immunization the mouse was sacrificed and adjuvant (Sigma, St. Louis, USA). Three days after homogenates were diluted 1:1 with Freund's incomplete RCC lesions obtained from 4 different patients. week intervals) with cell homogenates from primary An RBF mouse was immunized 5 times (with four

clones producing antibody binding to irrelevant

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Tissue samples taken from surgical specimens, and Scoring Procedure RESCRIATEY OF G 250

For immunofluorescent staining of RCC

to FITC (GAM-FITC) Nordic, Tilburg, The Netherlands) applied as a first step and goat antimouse Ig coupled

RAM-HPO was used as a second antibody and DAB/ $\mathrm{H}_2\mathrm{O}_2$ as

hybridoma culture fluid essentially according to van

blocks obtained from non adjacent parts, were tested.

autopsy and abortions were snap-frozen and stored at

Indirect immunoperoxidase staining of air-dried

sections, undiluted hybridoma culture fluid was

substrates. Sections were counterstained with

Muijen et al. Am. J. Pathol., 114 9-17 (1984).

and aceton fixed cryostat sections was done with

-70°C until used. For RCC, at least two tissue

hematoxylin.

Tissue Samples, Staining of Cryostat Sections

EXYMBLE 2

were subcloned and tested on other normal tissues. reacting with RCC and not with normal kidney tissue sections of RCC lesions and normal kidney. Clones medium from these clones was tested on crydstat picked and grown in suspension. Tissue culture diving spots on the RCC coated filter only were (DAB) and 0.03% H_2^{0} . Colonies producing antibodies washed and stained with 0.05% 3-3' diaminobenzidine conjugated to horseradish peroxidase (RAM-HPO), were removed and incubated with rabbit antimouse Ig antigens. After overnight incubation, the filters

as a second step. Sections were mounted and examined in a Leitz Orthoplan immunofluorescence microscope with Ploeaopak illuminator.

Cryostat sections of normal tissues were scored as negative when not a single cell was stained. In cryostat sections of benign and malignant tumors, percentages of cells stained per cm² were estimated visually. Four categories were arbitrarily distinguished. These were: negative, less than 1%, between 1% and 50%, and more than 50% of tumor cells stained.

EXYMBIE 3

Reactivity of Mab G 250-Staining of Cells obtained from Dr. L. J. Old, Memorial Sloane-Kettering Institute, New York, N. Y.

Single cell suspensions from fresh RCC specimen were obtained by collagenase treatment (Sigma, St.

Unfixed or acetone fixed cells grown on glass were examined for the presence of G 250 antigen by immunofluorescence. Undiluted culture medium was used as first step and GAM-FITC as second antibody. The cells were examined in a Leitz Orthoplan immuno-fluorescence microscope with Ploemopak.

RESULTS OF EXAMPLES 1-3

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Using the methods described above, a clone of hybridoma cells designated G 250 was isolated that produced antibody of the IgGl subclass reacting with

antibody concentration 40x and enhancing the staining Increasing the (50 cases), rejected transplant kidneys (8 cases) or embedded tissue sections of RCC. G 250 did not stain cryostat but not with formalin-fixed and paraffin

could be abolished by preabsorption with sup 2 of RCC The staining reaction of RCC cryostat sections staining of normal kidney structures. Сутосћет., 30, 491-493 (1980)), also did ndt lead to reaction with imidazole (Straus, J. Histochem. kidney biopsies from 5 SLE patients. any structure in cryostat sections of normal kidneys

Several other tumor types expressed the antigen Sepharose-G 250 columns. conjd not be purified by affinity chromatography on Proc. Natl. Acad. Sci., 76, 4350-4354, (1980), and ments done essentially according to Towbin et al., confd not be characterized in immunoblotting experiantigen present in crude RCC homogenates or in sup 2 suggests that G 250 recognizes a protein. The sensitivity to proteinase K staining reaction. tissue sections with 20mM NaIO did not reduce the digested sup 2 RCC fractions. Pretreatment of RCC but not with proteinase K (Sigma, St. Louis, USA)

RCC tumor cells always appeared to be cytoplas-The staining of positive non denerallly much lower. tumor cells and the intensity of staining were tumors stained, the percentages of G 250 positive detected by Mab G 250. However, the fraction of

matic.

In addition to the above-mentioned tumors, small numbers of a few other tumor types were tested (see Table 2). These were two Wilm's tumors (both negative), one prostatic carcinoma (negative), five adrenal cortical carcinomas (4 negative, one with more than 50% tumor cells positive), two liver cell carcinomas (both negative) and one carcinoma of the renal pelvis (more than 50% tumor cells positive).

Several benign tumors and premalignant lesions

were tested with Mab G 250 (see Table 2). In two renal adenomas (diameter 10mm and 20mm) all cell membranes were stained. Four cases of epitheliosis of the breast were negative, and ll cases of colonic cases with less than 50% of positive cells respectively. Of 6 mixed tumors of the salivary gland four tively. Of 6 mixed tumors of the salivary gland four tively. Of 6 mixed tumors of the salivary gland four tively. Of 6 mixed tumors of the salivary gland four tively. Of 6 mixed tumors of the salivary gland four tively.

Staining of Viable Cells and Cell Lines Using G 250 antibody, staining of cell membranes

was observed in unfixed single cell suspensions obtained after collagenase treatment of fresh RCC specimen. Also, membranous staining of RCC cellines SK-RC-1 and SK-RC-7 was observed while no reaction was seen with SK-RC-6. No fluorescence was seen on was seen with SK-RC-6. No fluorescence was seen on was seen with SK-RC-6.

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lines BSC-1 and CV-1 were both negative. transformed with adenovirus. The monkey renal cell

Enzyme Linked Immunosorbent Assay Using G 250 EXYMBIE 4

tant was stored at -70°C (= sup 1) and the pellet was After centrifugation (10 min 10,000g), the superna-Potter Elvehjem apparatus (5 strokes at 1500 rpm). homogenized in 3 ml of 10 mM Tris-HCl, pH 7.4, in a One gram of RCC or normal kidney tissue was

suspension was centrifuged for 10 min at 10,000g. taining 0.1% Triton X-100. After 10 min at 0°C this resuspended in 3 ml of 10mM Tris-HCl, pH 7.4, con-

Tris-HCl, pH 7.4 (= sup 2). The supernatant was dialyzed overnight against 10 mM

nM. As controls, Sp2/0 media, not containing H_2SO_4 and optical density readings were taken at 492 The reactions were stopped by adding 50 ul 2.5 M developed with 200 ul of 0-diphenylamine and H_2O_2 . with RAM-HPO, the plates were washed again and were 250 culture medium. After washing and incubating were incubated for 2 hours at 37°C with 100 ul of G ovalbumine in phosphate buffered saline, the wells After blocking remaining binding sites with 3% The contents were allowed to evaporate overnight at lutions of sup 1 and sup 2 in 0.1 M Na₂CO₃, pH 9.5. Teddington, U. K.) were filled with 100 ul of dishort, wells in microtiter plates (Sterilin Limited, homogenates was tested in a checkerboard assay. Presence of relevant antigen in the respective

antibody were used with and without RAM-HPO as second antibody.

Using normal kidney and RCC homogenates in ELISA as coating material, G 250 antigen was present only in the sup 2 fraction of RCC. The OD492 readings of wells coated with normal kidney extracts and incubated with G 250 culture medium did not exceed the values of Sp2/0 or RAM-HPO incubated wells. The values of Sp2/0 or RAM-HPO incubated wells were in the range of 0.01-0.03, whereas the OD492 readings on RCC sup 2 coated wells were in the range of from

EXYMBIE 2

Radioimmunoscintigraphy Using Mab G 250

G 250 antibody was purified from ascites fluid prepared in Fl (RBFxBalb/C) mice by chromatography on DEAE-AFFi-Gel Blue (Bio-Rad Laboratories, Richmond,

USA).

Purified G 250 IgGl was labeled with 99mTechnetium. Immunological activity of the 99mTc labeled G 250 preparation was evaluated in ELISA-tests.

Human RCC-xenografts were established by seeding 10⁶ SK-RC-1 subcutanously in nude mice. Tumor the tumor cells. As a control, a nude mouse bearing a tumor derived from human melanoma cell line BRO (Lockshin et al., <u>Cancer Res.</u> 45, 345-350, 1985) was

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a 30-40 min period 20 hours after injection. over a 5 min period 2 hours after injection and over puter system. 100,000 count images were acquired GCA 40A gammacamera, connected with an MDS A2 com-Scintigraphy was performed using a Gamma Toshiba

and 7 mm and one mouse bearing a melanoma tumor with Two mice bearing RCC tumors with diameters of b ground subtraction techniques were not utilized.

visualized weighed 60 mg and measured 5x5x4 mm. hours after injection (Fig. 3c). The smallest tumor bearing mouse, no tumor was visible after 2 or 20 In the melanoma accumulated in the RCC tumors. After 20 hours, 7% of the total body counts were injection as was the region of the liver (Fig. 3b). xenografts were distinctly visible 20 hours after tion of the RCC tumor could be made (Fig. 3a). The similar. After two hours, a scintigraphic distinc-G 250 peparations were evaluated in ELISA and were logical activities of the 99mTc labeled and unlabeled cific activity of 132 uC/ug protein. The immunoof 1.5 ug^{99m} Tc labeled G 250 antibody with a spea diameter of 6 mm were given intravenous injections

TOT RC 38, RC 3, RC69 and RC 154 Preparation of Hybridoma Cell Lines EXYMPLE 6

Source of Tissues

раемогградіс агеая мете taken from surgical specimens Tumor tissue samples, excluding necrotic and

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adenomas were obtained at autopsy. Renal several non adjacent tumor samples were taken. responding metastases were obtained. From each RCC In these two cases, primary tumor and cor-In two cases of RCC, autopsy material was primary and metastatic tumors were obtained at of various malignancies. In three cases of RCC, both

gland and skin. brain, lymph node, uterus, thyroid gland, adrenal prostate, lung, liver, breast, skeletal muscle, stomach, jejunum, colon, testis, cervix, pancreas, These included kidney, ureter, bladder, specimens. after death or from uninvolved parts of surgical Mabs from autopsies performed within a few hours Normal tissues used for specificity tests of

normal tissues obtained from different patients. three different tissue sections of the aforementioned All Mabs described here were tested on at least

Fetal kidney tissues of 11-, 13-, 14-, 15-, 18,

All tissues were snap-frozen and stored at -70°C measurements were obtained from abortions. and 20 week gestation as estimated from bodylength

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Preparation of Cell Homogenates

saline pH 7.4 (PBS). After centrifugation for 10 at 1500 rpm) in three volumes of phosphate buffered nized with a Potter Elvehjem homogenizer (5 strokes Tissue was homogecortex or medulla and from RCC. Cell homogenates were prepared from adult renal

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min. at 1000g, the supernatant was used for immunization purposes or for coating nitrocellulose fil-

Immunization

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Balb/c mice were used for immunization purposes. Each animal was immunized at least three times with homogenates prepared as described above. For the immunization with RCC, homogenates from three different patients were used. For the first injection of Freunds complete adjuvant (Sigma, St. Louis, USA). Each animal was injected with 0.5 ml of the mixture of tissue homogenate and Freunds adjuvant. Booster injections with Freunds incomplete adjuvant. Booster injections with Freunds incomplete adjuvant (1:1, sigma, St. Louis, USA) were given at two week intervalse. Three days after the last injection the mice vals. Three days after the last injection the mice sigma, St. Louis, USA) were given at two week intervalse. Three days after the last injection the mice vals.

Fusion of Spleen Cells with Sp2/0 and Detection of Hybridomas Producing Relevant Antibody

The spleen cells of immunized mice were fused

with Sp2/0 cells essentially according to Kohler and Milstein. After fusion, the cells were plated into 20 petridishes, diameter 5 cm, in soft agar (0.4%) and incubated for 10 days at 37°C in a CO₂ incubator. Then the agar was overlaid with nitrocellulose

the filters coated with the relevant cell homogenate Colonies producing antibodies that reacted with Sharon et al, Pro Watl Acad Sci, 76 1420-1424 (1979). The procedure followed was adapted from di-amino-benzidine and 0.03% $\rm H_{2}\rm O_{2}$ in 50 mM Tris-HCl, Netherlands), the filters were developed with 0.05% and 0.5% sarcosyl NL 30 (Ciba-Geigy B.V., Arnhem, The HCl, pH 7.4, containing 0.02% sodium-dodecyl-sulphate peroxidase. After extensive washing in lomM Trisrabbit-anti-mouse Ig conjugated to horseradish filters were removed and incubated for one hour with After overnight incubation on the soft agar, the calf serum to block remaining protein binding sites. filters were soaked in HAT-medium containing fetal Thereafter the filters were washed in sterile water. To reduce toxicity of the filters, the (30 Watt Philips TUV at 90 cm, 2 times 20 minutes the filters were air-dried and sterilized by UV light $_{
m TO}$ mJ of the cell homogenate was sucked through and on a sintered glass funnel of diameter 47 mm. Then were soaked in Hanks balanced salt solution and put The filters balanced salt solution and sonified. Cell homogenates were diluted 20 fold in Hanks These filters were prepared as follows: kidney relevant and irrelevant antibody producing human liver, as a first screen to discrimate between rated with a homogenate made from a normal adult dissolved in Hanks balanced salt solution or satuand a second filter, saturated with 0.75% gelatin responding to the homogenate used for immunization illers saturated with a cell homogenate cor-

but not with the filters coated with liver cell homogenate or gelatine were picked and grown in suspension in microliter plates. Undiluted culture media of these cells were tested on cryostat sections of several tissues. Colonies producing antibodies positive on adult renal tissue sections or RCC sections and negative on liver and lung tissue sections were subcloned and further analyzed on frozen sections were subcloned and further analyzed on frozen sections of other tissues.

EXYMBIE 1

Staining and Scoring Procedure of Mab RC 38,

To test the specificity of the Mabs and to

identify the structures stained, indirect immunoperoxidase staining was performed on frozen sections of various normal tissues, fetal kidneys and tumore as described by van Muijen et al., Am. J. Path., 114, 9-17 (1984), with the exception that 3,3'-di-aminobenzidine was used as substrate. Sections were counterstained with hematoxylin.

counterstained with hematoxylin.
Sections of normal tissues were scored negative

when not a single cell was stained. Tumors were stained, tumor sections in which only blood vessels were stained, tumor sections in which only blood vessels were stained were considered to be negative.

The subsite of the nephron stained by the Mabs was established by using generally accepted morpholowic criteria such as presence of brush-border and width of tubule lumen. The position of the tissue

section in the kidney was also taken into consideration. Rabbit-anti-human Tamm Horsfall protein (RAH-THP) was used to identify the sacending limb of Henle's loop and the distal convoluted tubule.

Double immunofluorescence staining was performed to identify any overlap of THP containing cells and the colls stained with the Mabs.

EXAMPLE 8

Test Set Of Poorly Differentiated Malignant Tumore

To confirm the diagnostic potential of Mab RC

To confirm the diagnostic potential of Mab RC

38, a test set of diagnostically difficult tumors was selected. Tumors of this test set included three cytological aspirates which present additional diagnostic difficulties, and poorly differentiated and malignant tumors with a histological appearance and sibility of RCC had therefore been considered by the pathologist in the differential diagnosis. The histological appearance of these cases included histological appearance of these cases included adenocarcinoma with clear cell, cribriform or acinar adenocarcinoma with clear cell, cribriform or acinar patterns, undifferentiated large-cell malignant tumors and spindle cell malignant tumors and spindle cell malignant tumors of these cases were included in the series of tumors used to test the specificity of the Mabs.

Tumors of the test set were stained and scored

Tumors of the test set were stained and scored by E. O. without knowledge of the final diagnosis.

KERDITE OF EXAMPLES 6-8

Mab RC 38, subclass IgGl, was derived from Origin and Subclass of Monoclonal Antibodies

from fusion of spleen cells of a mouse immunized. subclass IgG2b and IgG2a respectively were derived adult renal cortex homogenates. RC 69 and RC 154 with spleen cells from a mouse immunized with normal RCC homogenates. RC 3, subclass IgGl, was developed fusion of the spleen cells of a mouse immunized with

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All Mabs are only applicable on cryostat sec-

staining reaction was observed. When tested on formalin-fixed tissue, no

Staining of Normal Human and Fetal Tissue Sections

RC 38

by double immunofluorescence. present in the distal tubules--was seen as indicated plasm. No staining of the THP containing cells-tubular cells was mainly localized within the cytoing part of Henle's loop. Staining of the proximal cells of the proximal tubules up to the thin descendthe glomerular visceral epithelium and the epithelial In tissue sections of adult kidney RC 38 stained

thelial cells at the capillary loop stage and the stained differentiating visceral glomerular epi-In sections of the fetal metanephros, RC 38

redrous. most broximal part of the tubules connected to these

the mucous cells of the faveolar and glandular layer surface of the epithelium of the jejunum and colon, In non renal tissue sections, RC 38 stained the

negative (Table 3). lymph nodes were stained. Other tissues tested were addition sinusoidal lining cells in the liver and of the stomach mucosa and acini of sweat glands.

RC 3 and RC 69

In sections of the fetal metanephros the middle adjoining the outgoing tubule. ebithelial cells of Bowman's capsule in the region brush-border. RC 3 faintly stained the parietal proximal tubular cells was mainly associated with the parts of the nephron was observed. The staining of part of Henle's loop. No staining of more distal 69 stained the proximal tubules up to the descending In tissue sections of adult kidney RC 3 and RC

differentiating parietal epithelium were heavily stained with RC 69. Developing proximal tubules and eventually develop into the proximal tubule was limb of the S-shaped stage, the part that will

stained with both RC 3 and RC 69.

RC 3 or RC 69 (Table 5). All non renal tissues tested were negative with

In adult kidney tissue sections, a weak baso-BC J24

immunofluorescence, the distal tubular epithelium was tubules and small collecting ducts. In double cytoplasmatic basolateral staining of the distal was observed with RC 154 in addition to intense lateral staining of the proximal tubular epithelium

Staining of tumors

were negative (table 3).

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In the fetal metanephros essentially the same RC 124 ouly. proximal tubular epithelium was weakly positive with positive with both RC 154 and RAH-THP, whereas the

stained after the development of the capillary loop compared to the more proximal parts. Nephrons were distal parts of the differentiating nephron as distribution was seen: More intense staining of the

tissues sections was observed. Other tissues tested ducts in breast gland and follicles in thyroid gland In non renal tissue sections, weak staining of

RC 38 stained 95% of primary and 60% of metastatic from the urogenital tract are summarized in table 4. The staining results on RCC and other tumors

stain the sections of metastatic RCC tested. primary RCC, respectively. RC 69 and RC 154 did not RC 69 and RC 154 reacted with 70% and 40% of

(6x), lung (2x), ovary (3x) and soft tissue (1x). tumors included clear cell type tumors of the testis not stained by any of the Mabs (tables 4, 5). These tumors originating outside the urogenital tract were Other tumors of the urogenital tract and various

stained with the Mabs are indicated. The staining of In table 6 the percentages of tumor cells plood capillaries was often observed in all tumors. With RC 38 staining of endothelial cells of

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and tumor cells are stained. RC 3 and RC 69 (data not shown) the proximal tubules proximal tubules and tumor cells are stained. With With RC 38 a glomerulus, the of tumor sections. from a few positive tumor cells to diffuse staining RCC was heterogeneous for all four Mabs and ranged

The percentages of tumor cells stained were

quite similar in all pieces of one tumor studied

percentages of tumor cells were stained with the 1% of cells positive (see table 6) while higher seen with one or two Mabs corresponding to less than positive tumor cells per cm^2 of tumor section were In tissue sections of three RCC only 50-100 These percentages were estimated visualpercentages of tumor cells stained with the Mabs are In table 4 the observed in another tissue block. tissue block and a few positive tumor cells were except for two tumors where RC 3 failed to stain one

In primary RCC various combinations of the heterogeneity was seen as in primary RCC.

In metastatic RCC stained with RC 38 the same

possible combinations of antigen expression were not Note that some of the more of the Mabs, 7 with none. RCC tested with these three Mabs stained with two or 27 out of 41 primary expressed as shown in table 7. antigens recognized by RC 3, RC 69 and RC 154 are

ing metastatic lesion was available. RC 3 and RC 69 In 5 cases primary RCC as well as a correspond-

opserved.

other Mabs tested.

all five primary RCC and three metastases. none of the metastatic lesions, while RC 38 stained stained two and RC 154 one of these primary RCC and

Staining of test set tumors

other histochemical studies. found to be non-renal on the basis of clinical or tumors which were negative for RC 38 were eventually grounds, were negative with RC 38. All other 12 compatible with RCC on histological and clinical findings was RCC. Two other tumors which were, diagnosis based on additional clinicopathological body (table 8). In these 4 cases the ultimate test set of 18 tumors were stained with RC 38 anti-Four of the tumors (cases 1,2,3 and 4) in the

mm, no cells were stained by any of the Mabs. Fīve to stain any cell. In another adenoma, diameter 4 In one adenoma, diameter 10 mm, a few cells were Staining of renal adenomas

found to be negative. size from 2-4 mm were tested with RC 38 only and were other adenomas -four from one patient- ranging in stained with RC 38 while RC 3, RC 3 and RC 154 failed

specifically described reactants and/or operating similar success by substituting the generically or The preceding examples can be repeated with

preceding examples. conditions of this invention for those used in the

EXAMPLE 9 Radioimmunoscintigraphy and Biodistribution

in Nude Mice

tection of RCC. RC 38 and G 250 are useful RIS agents for the deground. In conclusion our studies indicate that resulting in excellent scans with very low backliver, spleen and kidneys had markedly improved III_In-labeled Mabs the biodistribution indices for bodies. Seven days after administration of the was possible with F(ab')2 fragments of both anti-Visualization of the tumor spleen with "Tc-label. ratios greater than 10 except the liver, kidneys and all three radionuclides all organs had tumor:tissue 24 hours after administration of the antibody. that maximum levels in the tumor were reached about Biodistribution indices (tumor:tissue ratios) showed clearly seen from 1,5-4 hours post injection. melanoma xenograft was observed. RCC could be the RCC xenograft or RC 38 and G 250 antibody in the No accumulation of an irrelvant antibody in showed tumor specific localization in the RCC xeno-Both antibodies sacrificed to study biodistribution. Various times after injection mice were imaged and globulins (Ig) and 99m Tc-labeled F(ab')2 fragments. using 99mTc, IS5 and III and I abeled intact immunosubcutaneous RCC and/or melanoma. Studies were done mmunoscintigraphic (RIS) agents in nude mice bearing Antibodies RC 38 and G 250 were used as radioiAt that time

Immunoscintigraphy of Tumor Bearing Human Kidney EXAMPLE 10

normal kidney tissue. No accumulation occured in supply of the tumor was poor as compared to the unexpected as first images showed that the blood .Yboditns 0250 ormee Slow accumulation was not labeled G 250. The first tumor slowly accumulated tumor bearing kidneys that were perfused with Ex vivo experiments were performed with two

normal kidney tissue.

side of the kidney coincided with tumor tissue cided with tumor tissue. A hot spot on the right tion by the pathologist all hot spots visible coinantibody, this ratio increased to 9:1. After examinahours with fresh Collins fluid to remove unbound G250 the tumor:kidney ration was 4:1. After washing for 7

administration of the labeled antibody.

the kidney.

An image was obtained l6 hours after the first

radiolabel was observed in the normal kidney tissue. radiolabel was observed, whereas no accumulation of

this kidney tumor was not an RCC but an oncocytoma of After examination by the pathologist it appeared that G 250 antibody. The tumor: kidney ration was 1,8:1. with perfusion fluid not containing radiolabeled

final image was obtained after washing for 10 hours that time the tumor: normal kidney ratio was 1:1,4.

An image was obtained 2 hours after the first

administration of 99 Tc labeled G 250 antibody.

In second tumor, fast accumulation of

protruding into the vena renalis. The tumor was diagnosed as a RCC.

CLAIMS

- Aonoclonal antibody specific to an antigen of renal cell carcinoma selected from the group consisting of antigen G 250, antigen RC 38, antigen RC 3, antigen RC 154.
- S. An antigen binding fragment of the monoclonal antibody of Claim 1.
- A composition for immunoscintigraphy of renal cell carcinoma, comprising a radiolabeled monotrom the group consisting of antigen G 250, antigen RC 3, antigen RC 69, and antigen RC 3, antigen RC 69, and
- The composition of Claim 3, wherein the antibody is apecific for antigen G 250 or RC 38.
- 5. A composition of Claim 3, wherein the antibody is radiolabeled with a radioisotope selected from the group consisting of ¹²³Iodine, ^{99m}Technetium or ¹¹¹Indium.
- A composition for immunoscintigraphy of RCC, comprising a radiolabeled antigen binding fragment of a monoclonal antibody specific for

an antigen selected from the group consisting of antigen G 250, antigen RC 3, antigen RC 38, antigen RC 69, and antigen RC 154.

- 7. The composition for immunoscintigraphy of RCC of Claim 6 wherein the antigen binding fragment is an F(ab')₂ or Fab' fragment.
- 8. A composition of Claim 6, wherein the antibody is specific for antigen G 250 or RC 38.
- 9. A composition of Claim 6, wherein the antibody or fragment is radiolabeled with a radioisotope 125 lodine, 99m the group consisting of 121 lodine, 99m technetium or 111 lodine.
- comprising a mixture of radiolabeled anti-RCC monoclonal antibodies or antigen binding fragments thereof, the monoclonal antibodies being antibodies specific for antigen G 250, antigen RC 38, antigen RC 3, antigen RC 59, and antigen RC 154.
- are F(ab') cor Fab' fragments.

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comprises antibody specific for antigen G 250 A composition of Claim 10, wherein the mixture

A composition of Claim 10, wherein the mixture and antibody specific for RC 38.

fragment of antibody specific for G250. pody specific for RC38 and an F(ab') or Fab' comprises an F(ab') or Fab' fragment of anti-

Isl lodine, 99m Technetium or Lil Indium. from the group consisting of 123 lodine, 125 Iois radiolabeled with a radioisotope selected A composition of Claim 10, wherein the antibody

carcinoma (RCC), comprising the steps of: 15. A method for detecting and localizing renal cell

RC 38, antigen RC 3, antigen RC 69, and group consisting of antigen G 250, antigen for an antigen of RCC selected from the an antibody or antibody fragment specific injecting a human subject parenterally with

(q with a signal generating label; antigen RC 154, the antibody being labelled

(၁ detecting the signal with a signal RCC: sug. antibody to accumulate at the site of the allowing sufficient time for the labeled

converting the detected signal to an image (p. detecting means; and

of the RCC.

- 16. A method of Claim 15, wherein the antibody tragment is an F(ab')₂ fragment or a Fab' tragment.
- L7. A method of Claim 15, wherein the antibody or tragment thereof is specific for antigen G 250 or RC 38.
- As. A method of Claim 15, wherein a mixture of antibody against RC 38 and G 250 or fragments thereof is injected.
- 19. A method of Claim 15, wherein the antibody is radiolabeled with a radioisotope selected from the group consisting of \$^{123}Iodine\$, \$^{131}Iodine\$, \$^{9m}Technetium or \$^{111}Indium\$.
- 20. A method of Claim 18, wherein the signal detecting means is a gamma camera.
- a tissue sample, comprising the presence of RCC in a tissue sample, comprising the steps of:

 a. contacting a tissue specimen from a patient suspected of having RCC with an antibody group consisting of G 250, RC 38, RC 3, RC 69 and RC 154 under conditions which RC 69 and RC 6

-05-

- b. determining whether the antibody binds to cells by immunohistochemical techniques, the binding of the antibody being an indicator of the presence of RCC.
- 22. A method of Claim 21 wherein the antibody is specific for antigen G 250 or RC 38.
- 23. A method of immunotherapy of RCC, comprising administering an anti-tumor amount of antibody specific for the RCC antigens selected from the group consisting of G 250, RC 38, RC 3, RC 69 and RC 154.
- 24. A method of Claim 23, wherein the antibody is
- conjugated to an anti-cancer agent.
- 25. A method of Claim 23, wherein a mixture of antibodies is administered.
- 26. A method of Claim 23, wherein the RCC antigens are selected from G 250 and RC 38.
- 27. A method of Claim 23, wherein the antibody is of the IgE class.
- 28. An antibody for targeting a cytotoxic cell to RCC, having dual specificity, a first specificity for an RCC antigen selected from the

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group consisting of RC 38, RC 4 and G250; and a second specificity for a human cytotoxic cell.

- 29. An antibody of Claim 28, wherein the human cytotoxic cell is a monocyte, a cytotoxic T cell or an MK cell.
- so. An antibody having a dual specificity, a first specificity for the T3 antigen of T cells and a second specificity for an RCC antigen selected from the group consisting of RC 4, RC 38 and G250
- 31. Monoclonal antibody RC 38.
- 32. Monoclonal antibody G 250.
- 33. Monoclonal antibody specific for G 250 or RC 38 of the IgE class.

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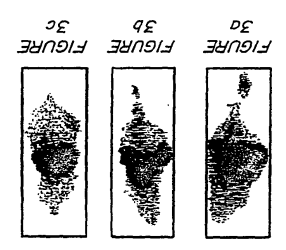
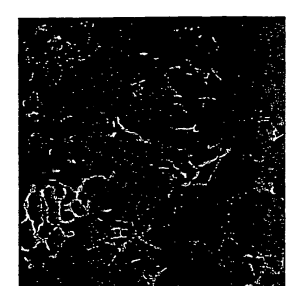


FIGURE 1





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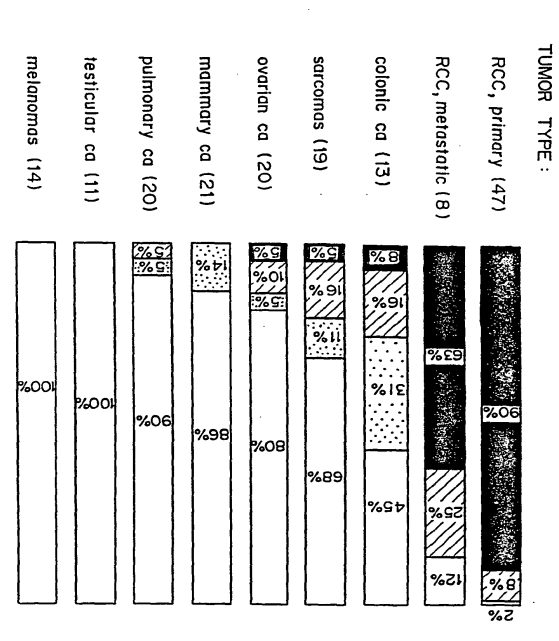


FIGURE 20

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FIGURE 4

Reactivity Pattern of Mab G 250



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4. As all searchable claims could be searched without effort justifying an additional fee, the intermational Searching Authority did not invite payment of any additional fee.
. 2. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers:
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V T OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE!
FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET PCT/ISA/210 (2)

ON INTERNATIONAL PATENT APPLICATION NO. US 8801511

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